Domain Structure of the *Staphylococcus aureus* Collagen Adhesin†


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ABSTRACT: Sequence analysis of surface proteins from Gram-positive bacteria indicates a composite organization consisting of unique and repeated segments. Thus, these proteins may contain discrete domains that could fold independently. In this paper, we have used a panel of biophysical methods, including gel permeation chromatography, analytical ultracentrifugation, circular dichroism, and fluorescence spectroscopy, to analyze the structural organization of the *Staphylococcus aureus* collagen adhesin, CNA. Our results indicate that the structure, function, and folding of the ligand-binding domain (A) are not affected by the presence or absence of the other major domain (B). In addition, little or no interaction is observed between the nearly identical repeat units within the B domain. We propose that CNA is indeed a mosaic protein in which the different domains previously indicated by sequence analysis operate independently.

Bacterial adherence to the host tissue is the first critical step in the pathogenic process of most infections. Organisms that cause infections in the extracellular space colonize the extracellular matrix. This adherence is mediated by a subfamily of bacterial surface proteins called MSCRAMMs (I–3). During the past decade several MSCRAMMs on Gram-positive bacteria have been identified and partially characterized (for reviews, see references 4–6). Sequence analyses of the MSCRAMMs revealed a common structural theme composed of unique and repeated sequences, and a multidomain organization of these proteins was therefore proposed. One such MSCRAMM, the collagen adhesin, CNA (Figure 1a), is present on some strains of *Staphylococcus aureus* and is composed of a N-terminal signal sequence followed by a 55 kDa segment of unique sequence called the A domain. This domain is followed by the B domain, which is composed of a 23 kDa subunit that is repeated 1–4 times in CNA from different *S. aureus* strains (7). The C-terminal portion of the protein consists of a cell wall attachment region, that contains the cell wall anchoring site LPKTG, a hydrophobic transmembrane region and a short cytoplasmic tail composed primarily of positively charged residues. It is believed that the protein is cleaved at the cell wall anchoring site and becomes covalently linked to the cell wall (8).

CNA mediates bacterial adherence to collagenous tissues such as cartilage, a process that is important in the pathogenesis of septic arthritis caused by *Staphylococci*. An isogenic mutant in which the *cna* gene has been inactivated is considerably less virulent than the wild-type CNA-expressing bacteria in a mouse model of septic arthritis (9). Furthermore, a *S. aureus* strain that normally lacks the *cna* gene has been inactivated by mutagenesis and is considerably less virulent than the wild-type strain (10).

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† Abbreviations: MSCRAMMs, microbial surface components recognizing adhesive matrix molecules; CNA, *Staphylococcus aureus* collagen adhesin; SDS-PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis; EDTA, ethylenediaminetetraacetic acid; ESI/MS, electrospray ionization mass spectrometry; UV/vis, ultraviolet/visible; CD, circular dichroism; SPR, surface plasmon resonance spectroscopy; HEPES, N-(2-hydroxyethyl)piperazine-N′-2-ethanesulfonic acid; BSA, bovine serum albumin; amino acids are represented by standard one-letter codes.

MRGSHHHHHHGS is the amino acid sequence of the unstructured His6-tag required for purification.

Figure 1: (a) Schematic of the *S. aureus* FDA 574 collagen-binding MSCRAMM, CNA. The putative signal peptide (S), nonrepetitive binding domain (A), domain of three subunits (B1B2B3), cell wall domain (W), membrane spanning domain (M), and charged carboxyl-terminal domain (C) are indicated. (b) Recombinant CNA proteins used in this study that mimic portions of the collagen-binding MSCRAMM, with the inclusive residues indicated. The A domain subfragment (M19) for which the crystal structure has been solved spans residues 151–318 (11).
gene becomes more virulent in this model when the cna gene is introduced (9). Recent studies suggest that the recombinant CNA A domain, when used as a vaccine, provides protection in a mouse model of staphylococcal-induced sepsis (10).

The collagen-binding function of CNA has been located to the A domain, and a recombinant protein containing a 19 kDa subfragment of the A domain has collagen-binding activity (11), although a recombinant version of the full-length A domain appears to bind collagen more effectively than the shorter subfragment (12). We recently solved the crystal structure of the 19 kDa collagen-binding subfragment (13). This polypeptide folds as a jellyroll composed of two $\beta$-sheets connected by two short $\alpha$-helices. One of the $\beta$-sheets has a noticeable “trench” transversing it. Molecular modeling studies showed that this trench can accommodate a collagen triple helix, and analyses of single point mutations in the trench suggest this does in fact represent the collagen-binding site and several residues making contact along the extended collagen macromolecule. Such a binding scheme has been suggested recently for other collagen-binding proteins (14, 15).

The B domain of CNA in S. aureus strain FDA 574 is a triplet of repeat units. The first and third repeat units of the B domain are sequentially identical; the second varies by a triplet of repeat units. The first and third repeat units of the proteins (14, 15) are identical to the now-determined sequence except for the following: Arg399→Ala, Ile665→Thr, and Ile852→Thr.

**Bacterial Growth Conditions.** Recombinant E. coli (JM101) were grown at 37 °C in Luria Broth (Life Technologies) containing 100 μg/mL ampicillin. Overnight cultures of stationary-phase bacteria were used to inoculate fresh media (1:1000), and the cells were allowed to grow until OD$_{600nm}$ = 0.6–0.8 (approximately 6 h). Protein expression was induced by addition of isopropyl-β-D-galactoside to a final concentration of 0.2 mM, and the culture was incubated for an additional 3 h. Bacteria were collected by centrifugation and resuspended in a minimal volume of 5 mM imidazole, 100 mM NaCl, 4 mM Tris-HCl, pH 7.9, and frozen at –70 °C.

**Isolation of Recombinant Proteins.** Cell pellets were thawed, passed through a French press (11 000 lb/in$^2$) 3 times to lyse the cells, and centrifuged at 14 krpm for 20 min to remove cell debris and insoluble proteins. The supernatants were filtered through a 0.45 μm membrane, and recombinant proteins were purified using metal-chelating affinity chromatography. Bacterial lysates were applied to a HR10/10 (Pharmacia) column of nickel-charged iminodiacetic acid Sepharose (Sigma), and bound protein was eluted with a 200 mM linear gradient of 0–200 mM imidazole in 4 mM Tris-HCl, 100 mM NaCl, pH 7.9. In general, this purification step yielded proteins that were >95% pure as estimated by SDS–PAGE. The recombinant AB$_1$ preparation, however, required an additional purification step. Fractions containing

| Table 1: Synthetic Oligonucleotides Used in Plasmid Constructions |
|---------------------|------------------|------------------|
| construct | base pairs | amino acids | oligonucleotide$^a$ |
| A | 238–1744 | 30–531$^b$ | $\text{BamHI primer \#5: 5'}$ |
| AB$_1$ | 238–2314 | 30–721 | $\text{P3 primer: 5'}-\text{CGGGATCCTGAGGTTATATGATGAAACC-3'}$ |
| B$_1$ | 1753–2314 | 535–721 | $\text{BamHI primer \#5: 5'}-\text{CGGGATCCCGGAAATATAGTTA-3'}$ |
| B$_1$B$_2$ | 1753–2875 | 535–908 | $\text{B1 (3') primer: 5'}-\text{CGGGATCCCGGAAATATAGTTA-3'}$ |

$^a$ Restriction endonuclease sites are in boldface type. $^b$ Not including N-terminal histidine tag.
recombinant AB₁ (as determined by SDS–PAGE) were pooled and dialyzed against 20 mM tris-HCl, pH 8.0, before being applied to a HR10/10 Q Sepharose column (Pharmacia). Bound protein was then eluted using a 100 mM NaCl gradient of 0–0.5 M NaCl, 20 mM Tris–HCl, pH 8.0. During all chromatographic procedures, the eluate was monitored by absorbance at 280 nm and collected fractions were examined for purity by SDS–PAGE. Fractions containing only the desired product were pooled and concentrated using an Amincon ultrafiltration system. Concentrated proteins were then dialyzed extensively against 10 mM Na₂HPO₄, 100 mM NaCl, pH 7.0, fast-frozen, and kept at −70 °C until use.

**Verification of Protein Identity.** To verify the protein’s identities, electrospray mass spectrometry measurements of each were performed at the University of Texas–Houston Health Science Center. Samples were dialyzed 3 × 1 L against 50 mM EDTA in water to remove Ni²⁺, and then dialyzed 3 × 1 L against water to remove excess EDTA. All aqueous stock sample solutions were diluted with ESI/MS solvent, consisting of 100/100/10 (v/v/v) methanol/water/glacial acetic acid. Instrument setup and calibration were performed using a mixture of horse skeletal muscle apomyoglobin and BSA (2.5 pmol/L each) in ESI/MS solvent. Analyses were performed using a Finnigan TSQ70 (upgraded to TSQ700) mass spectrometer fitted with a modified Vestec ESI source. Duplicate spectra were acquired for each sample. The instrument was calibrated before each run using a series of standards; deviation between theoretical and measured molecular masses of the standards was consistently less than 0.02%. Table 2 lists the molecular mass of each recombinant protein as determined from the amino acid sequence, measured by mass spectrometry, and calculated from sedimentation velocity parameters.

**Gel Permeation Chromatography.** Effective radii were measured at room temperature using gel-permeation chromatography. Protein samples were diluted in 10 mM Na₂HPO₄, 100 mM NaCl, pH 7.0, applied to a HR10/30 Superdex 200 column, and eluted with the same buffer at a flow rate of 0.5 mL/min. The column was calibrated using a standard globular protein kit (Pharmacia). Radii were then calculated using sample elution volumes and standard curves as described in the calibration kit.

**Sedimentation Velocity Measurements.** Prior to analysis, protein solutions were dialyzed 3 × 1 L against 10 mM Na₂HPO₄, pH 7.0, to remove NaCl that may damage the instrument’s centerpieces. Experiments were performed using a Beckman XL-A analytical ultracentrifuge, equipped with an AN-60 Ti four-position rotor and absorption optics. All samples were centrifuged at 20 °C and 60 krpm in double-sector aluminum centerpieces. Sedimentation data were analyzed with the UltraScan software package² using the van Holde–Weischet (21), second moment (22), and finite element analysis (23, 24) methods. The van Holde–Weischet method was used to confirm sample homogeneity and identify the possible concentration dependence of the sedimentation coefficient, s. The second moment method was used to ensure that the scans included in the analysis did not exhibit time-dependent variations such as aggregation, degradation, or pelleting. Finally, to determine the sedimentation and diffusion coefficients, the experimental data were fit directly to a finite element solution of the Lamm equation as described by Demeler and Saber (25). The partial specific volume, \( \bar{\rho} \), of each protein was estimated from the amino acid sequence by the method of Sober (26) and was temperature-corrected. To model the overall geometry of the molecules, axial ratios were estimated for prolate and oblate ellipsoids from the experimentally determined sedimentation and diffusion coefficients as described in ref 27.

**Absorption Spectroscopy.** Absorption measurements were taken at ambient temperature (23 ± 2 °C) using a Beckman DU-70 UV/vis spectrophotometer and a 1.0 cm path length cuvette. All spectra were corrected for background signal, and the reported absorbance maxima are ±1.0 nm. Since the proteins in this study all contain chromophoric amino acids, the concentration of each protein solution was determined spectrophotometrically using the Edelhoch method of UV-light absorption (28). Extinction coefficients were experimentally determined as described by Gill and von Hippel (29), employing Pace et al.’s values for the extinction coefficients of the individual residues (30, 31).

**Steady-State Fluorescence Spectroscopy.** Fluorescence excitation and emission spectra were collected at ambient temperature using an Aminco SLM 4800C fluorimeter equipped with a 450 W xenon lamp, employing excitation and emission band-passes of 2 nm. The excitation source was set at 295 nm to preferentially select tryptophyl emission. Protein samples, in 10 mM Na₂HPO₄, 100 mM NaCl, pH 7.0, were stirred throughout each scan to prevent photo-bleaching. All scans were background-subtracted, and all reported maxima were reproducible within ±1.0 nm. The relative fluorescence quantum yields, \( \phi_f(x) \), were determined using the calculations outlined by Petrich and co-workers (32):

<table>
<thead>
<tr>
<th>Table 2: Summary of Physical Parameters</th>
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<tr>
<td>molecular mass (kDa)⁴⁶</td>
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<tr>
<td>species</td>
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<tr>
<td>A</td>
</tr>
<tr>
<td>AB₁</td>
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<tr>
<td>B</td>
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<td>B₁B₂</td>
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⁴ Values represent molecular mass calculated from the published sequence, mass spectrometry, and sedimentation velocity measurements, respectively. ³ Sedimentation coefficients obtained from the finite element analysis and calculated using the molecular weight determined by sedimentation velocity. ⁵ Assuming a hydrated sphere. The data shown here are an average of values obtained by gel-permeation chromatography and sedimentation velocity measurements. Radii expected for globular proteins of these molecular masses are shown in parentheses and calculated from formulas outlined by Uversky (46).
where OD(x) is the optical density of recombinant protein x at the excitation wavelength and \( I_{em}(x,v) \) is the emission intensity at the excitation wavelength. For these studies, fluorescence quantum yields were determined relative to that of CNA B1. Chaotropic protein denaturation was induced by equilibrating the samples overnight with 6.0 M guanidine hydrochloride (ultrapure grade, USBiochem), 10 mM Na2HPO4, 100 mM NaCl, pH 7.0, at ambient temperature. Emission spectra of guanidine hydrochloride in buffer were subtracted from the corresponding sample scans.

Time-Resolved Fluorescence Emission Anisotropy. Proteins were labeled with Texas Red Sulfonyl Chloride (Pierce Chemical) to measure rotational tumbling ratios. Each protein was diluted into 10 mM Na2HPO4, 100 mM NaCl, pH 9.0, to a final concentration of 0.2 mg/mL. The labeling was performed as previously described (33, 34), with the following modifications. One milligram of Texas Red (\( \lambda_{em, max} = 596 \text{ nm in buffered aqueous solution} \)) was added to the protein solution, which was then mixed gently and incubated at room temperature for 1 h. To remove excess dye, the protein solution was diazylated exhaustively against 10 mM Na2HPO4, 100 mM NaCl, pH 7.0.

Fluorescence lifetimes, \( K(t) \), and fluorescence anisotropy decays, \( r(t) \), were obtained using the time-correlated, single-photon counting technique \( (35, 36) \). A time scale of approximately 15 ns was used to collect the fluorescence decays. All measurements were performed at 20.3 °C and with \( \lambda_{ex} = 570 \text{ nm and } \lambda_{em} > 610 \text{ nm} \). The anisotropy decays of the proteins were fit to the function \( r(t) = r_1(0) \exp(-t/\tau_1) + r_2(0) \exp(-t/\tau_2) \), with the short time constant fixed at the value obtained for the probe alone. Calculations for the determination of the order parameter and hypothetical cone semiangle are described in ref. 32.

Circular Dichroism (CD). Far-UV circular dichroism data were collected using a Jasco J720 spectropolarimeter calibrated with d-10-camphorsulfonic acid, employing a bandpass of 1 nm and integrated for 4 s at 0.2 nm intervals. All sample concentrations were less than 15 \( \mu \text{M in 1 mM Na2HPO}_4 \), 10 mM NaCl, pH 7.0. Spectra were recorded at ambient temperature in cylindrical 0.5 mm path length cuvettes.

Twenty scans were averaged for each spectrum, the contribution from buffer was subtracted, and quantitation of secondary structural components was performed using software provided by N. J. Greenfield (UMDNJ—Robert Wood Johnson Medical School, Piscataway, NJ) and D. Greenwood (Softwood Co., Brookfield, CT). We initially used four deconvolution programs (SELCON, VARSCLC1, CONTIN, and CD estima) that were derived from databases of known protein structures (37–40). The reliability of these programs in predicting the secondary structure of CNA domains was assessed by analyzing the spectra obtained for M19, a truncated form (19 kDa, residues 151–318) of the A domain that encompasses the collagen-binding region. The structure of M19 has been solved using X-ray crystallography and consists of 8% \( \alpha \)-helix, 53% \( \beta \)-sheet, and 39% coil (13). The CONTIN analysis consistently yielded exaggerated \( \beta \)-sheet content, while the CD estima program dramatically underestimated the percentage of \( \beta \)-sheet structure. Averaging data from all four programs yielded the same results as comparison of SELCON and VARSCLC1 only, but with significant error (up to 50%). As a result, we have chosen to report an average of results obtained from the SELCON and VARSCLC1 analyses only. For M19, this yielded secondary structure percentages that are comparable to the data obtained from the crystallographic analysis: 12 ± 4% \( \alpha \)-helix, 49 ± 4% \( \beta \)-sheet, and 39 ± 13% other components.

Thermal Denaturation. Protein unfolding was induced by heat and monitored by far-UV CD. The thermal denaturation (ramp rate of 20 °C/h) of each protein in 1 mM Na2HPO4, 10 mM NaCl, pH 7.0, was followed at 208 nm by far-UV CD using the parameters described above. The reversibility of the thermal denaturation was determined immediately after the unfolding analyses, and the samples were cooled at a ramp rate of 20 °C/h.

Surface Plasmon Resonance Spectroscopy (SPR). Analyses were performed using the BIACore system. Bovine type I collagen (Collagen Corp., Fremont, CA) was immobilized on a CM5 sensor chip as described previously (11). Recombinant CNA proteins in 150 mM NaCl, 50 mM HEPES, 0.05% P-20 surfactant, pH 7.4, were flowed over three flow cells containing different amounts of immobilized collagen. All data were corrected for the response obtained using a flow cell that was not coated with collagen. Analytical conditions and data analysis were as described previously (12, 41). No mass transport effects were observed in these measurements.

ELISA Assays. Microtiter plates were coated with 10 \( \mu \text{g of type I collagen per well in 10 mM Na2HPO}_4 \), 100 mM NaCl, pH 7.0, overnight at 4 °C. Collagen-coated wells were then washed with the same buffer and incubated for 1 h in the presence of 5% BSA in 10 mM Na2HPO4, 100 mM NaCl, pH 7.0. Wells were again washed with buffer, and varying concentrations of recombinant CNA protein, in buffer containing 0.1% BSA, were added. After 3 h, unbound protein was removed by washing the wells with the same buffer. To detect bound protein, the wells were incubated with a monoclonal antibody that recognizes the histidine tag of the recombinant proteins (6xHis Monoclonal Antibody, Clontech), followed by alkaline phosphatase-conjugated goat anti-mouse IgG antibody. Relative binding was measured by monitoring the sample absorbance at 405 nm following the addition of p-nitrophenyl phosphate in 1 M diethanolamine, 0.5 mM MgCl2, pH 9.0.

RESULTS

Expression and Purification of Recombinant Proteins Mimicking Regions of CNA. A model of the collagen-binding MSCRAMM from S. aureus FDA 574 has been described previously, in which a multidomain nature of this protein was proposed (8). We have now analyzed recombinant proteins containing individual (A and B), homodimeric (B1B2), and heterodimeric (AB1) domains of the MSCRAMM with the goal to evaluate this model. SDS-PAGE, followed by staining with Coomassie Brilliant Blue, revealed that each preparation of recombinant protein migrated as a single band, but at unexpectedly high molecular masses due to their acidic natures (Figure 2). No degradation
The negative slope of the distribution of A and AB$_1$ indicates slight of 0.5 refers to the midpoint in the sedimentation velocity boundary. The data shown refer to diffusion, temperature, and buffer-corrected of concentrations and yielded a not indicate aggregation of the A domain over a wide range of concentrations using a Beckman XL-A analytical ultracentrifuge did similar mass (Table 2). Sedimentation velocity measure-
ments using a Beckman XL-A analytical ultracentrifuge did similar mass (Table 2). If we assume a prolate ellipsoid model for this protein, the calculated axial ratio would be 12.8. Hence, the recombinant CNA A domain recombinant is a primarily \( \beta \)-sheet protein that contains a small percentage \( \alpha \)-helix (Table 3). This composition of secondary structure elements is similar to that for M19, the CNA A domain truncate that encompasses the collagen-binding motif (11), for which the crystal structure was recently solved (13). Thermal dena-
turation, as followed by intensity of the far-UV CD spectrum, revealed that the A domain is quite thermostable and unfolds reversibly over a range of 30 °C (60–90 °C), undergoing a multiphasic thermal denaturation with a highly populated intermediate state apparent at approximately 74 °C (Figure 5).

The recombinant A domain protein exhibits an absorbance maximum of 278 nm and has an experimentally determined extinction coefficient of 58 740 M$^{-1}$ cm$^{-1}$. The fluorescence quantum yield of A is 15% of that for B$_1$, and the five tryptophans of the A domain reside in an overall nonpolar environment (\( \lambda_{em}^{max} = 330 \) nm when \( \lambda_{em} = 295 \) nm) as shown by steady-state emission spectroscopy. The A domain’s tryptophyl emission spectrum red-shifts upon denaturation in 6 M guanidine hydrochloride (\( \lambda_{em}^{max} = 352 \) nm) as all tryptophans become fully solvated.
for any single repeat unit of the B domain since the amino measurements of B1 revealed that this protein eluted earlier the residues are identical. Gel-permeation chromatography dimerization at high protein concentrations (the monomeric in solution (Figure 3), and attempts to induce velocity analyses demonstrated that a single repeat unit is globular protein of the same mass, Table 2). Sedimentation yielding an effective radius of 26.3 Å (versus 22.5 Å for a than would be expected for a protein of this molecular mass, unit is 278 nm, and the extinction coefficient is 37 070 M -1 cm -1 ).

Structure of a CNA B Domain Repeat Unit. We propose that the structural data obtained for the B1 subunit are valid for any single repeat unit of the B domain since the amino acid sequences of the repeat units are so similar (>99% of the residues are identical). Gel-permeation chromatography measurements of B1 revealed that this protein eluted earlier than would be expected for a protein of this molecular mass, yielding an effective radius of 26.3 Å (versus 22.5 Å for a globular protein of the same mass, Table 2). Sedimentation velocity analyses demonstrated that a single repeat unit is monomeric in solution (Figure 3), and attempts to induce dimerization at high protein concentrations (>200 μM) in order to mimic the B1B2 recombinant were unsuccessful. In addition, the sedimentation velocity measurements gave a F/F o ratio of 1.42. When modeled as a prolate ellipsoid, this would represent a shape with an axial ratio of 7.7. Together, the gel-permeation chromatography and sedimentation velocity results indicate that a CNA B domain repeat unit is nonglobular.

The far-UV CD spectrum of B1 contains a relative minimum at 214 nm, maxima at 203 and 228 nm, and significant negative molar ellipticity at 190 nm (Figure 4a). These characteristics are typical of a structure rich in β-sheet. Deconvolution of this spectrum yielded secondary structural content that resembles that of the A domain (Table 3). However, the arrangement of the secondary structural motifs in the two domains is clearly not the same since the far-UV CD spectra of the two proteins are quite different. Heat denaturation of the recombinant single B domain repeat unit followed by changes in the far-UV CD spectrum demonstrated that this single repeat unit is particularly stable with respect to heat, denaturing reversibly in the range 68–82 °C via an apparently single, sharp transition (Figure 5).

The absorbance maximum of the recombinant B1 protein is a dimer of two B repeat units covalently linked at the C-terminus of the first and the N-terminus of the second. Since the sequence of the second repeat unit of the CNA B domain mimics exactly that of the first except for one residue (and the third repeat unit is identical to the first), we suggest that this dimer, B1B2, is a model for any two covalently joined B domain repeat units. The far-UV CD spectrum of B1B2 overlays that obtained for B1 (Figure 4a), implying that the secondary structures of the B repeat units do not change dramatically when they are covalently linked in series. Deconvolution of the far-UV CD spectrum produced the same composition of structural components as for B1, as was expected (Table 3). In addition, the secondary structural elements in B1B2 denature upon heating apparently as they do in B1, for the completely reversible thermal denaturation of B1B2 occurs at 70–82 °C and is also a simple, two-state transition (Figure 5).

Gel-permeation chromatography measurements indicate that the effective radius of B1B2 is somewhat greater than that expected for a 44 kDa globular protein (31.3 vs 28.8 Å, Table 2). Sedimentation velocity experiments determined that neither degradation nor multimer formation of B1B2 occurs at high protein concentrations and the protein has a F/F o ratio of 1.29 (Figure 3, Table 2). B1B2, therefore, also is nonglobular, yet not to the extent that B1 is.

The absorption spectrum of the recombinant B1B2 heterodimer has a maximum of 278 nm, and the protein’s extinction coefficient is 74 450 M -1 cm -1 . The 10 tryptophans of B1B2 are in an environment comparable with those of B1 (λ max = 341 nm, λ max = 295 nm). B1B2’s fluorescence quantum yield is twice that of B1. In fact, a theoretical construction of the B1B2 emission spectrum shown in Figure 6a closely duplicated the measured spectrum of B1B2 (the theoretical construction of the emission spectrum from eq 3 is discussed in detail below for AB1). The emission maximum of B1B2 in 6 M guanidine hydrochloride shifts to 352 nm as all tryptophans become completely solvated upon protein denaturation.

We would expect that if no significant structural reorganization occurs among the B repeat units upon joining to form the dimer, B1B2 would have a significantly greater diameter than B1. One method to compare relative protein sizes employs time-resolved fluorescence anisotropy. This technique follows the rotation of molecules in solution. Since large proteins tumble slower than small ones, we would expect the anisotropy decay (and, therefore, the rotational rate) of B1B2 to be much longer lived than that of B1 if the individual repeats fold independently and are simply joined by a polypeptide tether sequence. For this analysis, B1 and B1B2 were labeled with the visible light fluorescent probe Texas Red, and emission greater than 610 nm was collected.

The fluorescence lifetime of the Texas Red probe fit well to a single exponential of 4.2 ns. The fluorescence lifetime of this label did not change dramatically upon attachment to the B1 (or B1B2) recombinant protein. The fluorescence lifetime decays of B1 and B1B2 both fit to an exponential of the form: $K(t) = A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2) = 0.80 \exp(-t/4.5 \text{ ns}) + 0.20 \exp(-t/1.0 \text{ ns})$, indicating that the local environment of the probe was similar in these two proteins.
The anisotropy decay of the Texas Red dye alone in solution fit well to a single exponential of 240 ps (1.38) (Figure 7a). If the data were fit to two exponentials, a long component \( r_1 (0) \) = 15.9 ns; \( r_2 (0) \) = 0.13, \( r_2 (0) \) = 0.01 was observed. This results in a hypothetical cone semiangle, \( \theta_0 \) (within which the transition dipole moment can diffuse), of approximately 90°, as was expected for the probe free in solution.

The fluorescence anisotropy decay of B1 tagged with Texas Red fit well to two exponentially decaying components: \( r(t) = r_1 (0) \exp(-t/\tau_{1}) + r_2 (0) \exp(-t/\tau_{2}) = 0.09 \exp(-t/0.24 \text{ ns}) + 0.12 \exp(-t/8.8 \text{ ns}) \), \( \chi^2 = 1.25 \) (Figure 7b). This indicates that we were probing the rapid librational motion of the probe with respect to the recombinant protein as well as the overall tumbling motion of the protein macromolecule itself. The rotational anisotropy data for B1B2 were also resolved into two components: \( r(t) = 0.06 \exp(-t/0.24 \text{ ns}) + 0.15 \exp(-t/11.9 \text{ ns}) \), \( \chi^2 = 1.26 \) (Figure 7c). The second component of the anisotropy decay (\( \tau_{2} \)) of B1B2 is substantially longer-lived than that of B1, as would be expected for a molecule of greater length. The hypothetical cone semiangle was determined to be approximately 34° for Texas Red attached to B1 and 27° for B1B2. The fluorescent probe Texas Red alone in solution; (b) the labeled single B domain repeat unit, B1; (c) the labeled B domain repeat unit dimer, B1B2. The residuals corresponding to emission polarized parallel and perpendicular to the excitation source are overlaid above each panel. All time-resolved measurements have an error of approximately 10%.

**FIGURE 6:** Comparison of the fluorescence emission spectra of recombinant CNA proteins. (a) B1 and B1B2. The dashed line is the construction of a B1B2 spectrum using the B1 spectrum, where B1B2(calc'd spectrum) = 2 × B1(meas'd spectrum). (b) A, B1, and AB1. The dashed line is the construction of the AB1 spectrum using eq 3. All sample concentrations were 1.0 μM, and λex = 295 nm.

**FIGURE 7:** Parallel and perpendicular fluorescence intensity profiles of the recombinant CNA proteins from which the anisotropy decay is calculated (λex = 570 nm, λem > 610 nm, 20.3 °C). (a) The fluorescent probe Texas Red alone in solution; (b) the labeled single B domain repeat unit, B1; (c) the labeled B domain repeat unit dimer, B1B2. The residuals corresponding to emission polarized parallel and perpendicular to the excitation source are overlaid above each panel. All time-resolved measurements have an error of approximately 10%.
indicative of a protein having significant \(\alpha\)-sheet character, with an intensity minimum at 214 nm and relative maxima at 200 and 229 nm. The composition of secondary structural components of AB1 is similar to that of the other constructs (Table 3), yet its profile (Figure 4b) does not superimpose that of the recombinant A or B domain shown in Figure 4a. However, the percentages and arrangements of \(\alpha\)-helix, \(\beta\)-sheet, and other components of A and B1 do not undergo any change upon forming AB1 when the far-UV CD spectrum may be constructed from the weighted sum of A and B1’s spectra that duplicates the observed AB1 spectrum in Figure 4b:

\[
[\theta]_{\text{AB1}}(\text{calc’d}) = \frac{\text{(residues)}_A([\theta])_A + \text{(residues)}_B([\theta])_B}{\text{(residues)}_{\text{AB1}}}
\]

where \text{residues} is the number of amino acids in each protein. This is indeed the case: the theoretical construction and measured curve are nearly superimposable (Figure 4b). Hence, the organization of secondary structural components in A and B1 is the same in the individual recombinants and the heterodimer.

The effective radius (42.8 Å) of AB1 is significantly greater than expected for a globular 78 kDa protein. This protein is monomeric by sedimentation velocity analysis (Figure 3), and its \(F/F_o\) ratio of 1.56 is intermediate between those of A and B1, indicating a shape less globular than B1, but more so than A alone.

The thermal unfolding of AB1 is complex: no obvious intermediate state is observed, and the transition from native to denatured protein is diffuse, with a \(T_m\) of 70 °C and a dynamic range of more than 25 °C (Figure 5). This would indicate a denaturation via an unresolved intermediate state. Unlike the unfolding of the other recombinants, the denaturation of AB1 is not completely reversible: only approximately 70% of the original AB1 conformation is regained upon cooling.

The absorbance maximum of the recombinant AB1 heterodimer is 278 nm, and its extinction coefficient is 92 750 M\(^{-1}\) cm\(^{-1}\). From the steady-state emission spectrum of AB1, we conclude that its 10 tryptophans reside in an overall environment of intermediate polarity (\(\lambda_{em}^{\text{calc’d}} = 337\) nm, \(\lambda_{em} = 295\) nm) in relation to the other 3 recombinant proteins in this study. The fluorescence quantum yield of AB1 is 0.85 with respect to B1. As is expected, the fluorescence emission spectrum of AB1 in 6 M guanadine hydrochloride indicates all 10 tryptophans become fully solvated upon denaturation.

A theoretical construction similar to that described above for the secondary structure of AB1 (eq 2) may be used to determine the change, if any, in tertiary structure upon forming the mixed-domain protein. A theoretical fluorescence emission spectrum of (A + B1) may be constructed based on the relative quantum yields (A has a fluorescence quantum yield of 0.15 with respect to B1) and spectra shape and positions of the individual A and B1 units:

\[
I_{\text{AB1}}(\text{calc’d}) = \frac{(\phi_F^{\text{rel}})_{A}(I_A) + (\phi_F^{\text{rel}})_{B1}(I_{B1})}{(\phi_F^{\text{rel}})_{A} + (\phi_F^{\text{rel}})_{B1}}
\]

where \(I_{\text{AB1}}(\text{calc’d})\) is the calculated emission spectrum of AB1. \((\phi_F^{\text{rel}})_{A}\) is the quantum yield of A relative to B1, and \(I_A\) is the emission intensity of the A domain protein. If the tryptophan environments of both A and B1 are not altered upon complexation (for example, surface-exposed tryptophans of the A and B1 proteins becoming buried in the A–B1 interface), this calculated spectrum should duplicate the observed AB1 spectrum in Figure 6b. The observed and calculated spectra are nearly identical, providing additional evidence that (1) the local environments about the tryptophans of the A and B1 proteins becoming buried in the A–B1 interface, and (2) no tryptophans reside at the A–B1 interface.

Collagen Binding Is Not Influenced by the B Domain. An ELISA demonstrating the four CNA recombinant proteins binding to immobilized type I collagen is shown in Figure 8. Neither B1 nor B1B2 exhibited collagen-binding activity. In this assay, the affinities of A and AB1 for collagen were comparable. Figure 9 illustrates the normalized SPR responses of A and AB1 flowed over a BIAcore sensor chip coated with type I collagen (the responses of B1 and B1B2 were undetectable). It is apparent from these SPR profiles
that addition of the B₁ repeat unit to the A domain does not significantly affect the association and dissociation rates of the protein/collagen interaction. In addition, the recombinant A and AB₁ proteins compete for sites within the collagen macromolecule. Figure 10 demonstrates the inhibition of AB₁ by biotinylated A, in which the collagen binding of AB₁ was completely blocked by A. The reverse experiment yielded similar results: the inhibition of A by AB₁ is complete (data not shown). Data in Figures 8–10 illustrate that the presence of a B domain repeat unit does not alter the collagen-binding characteristics of the A domain.

DISCUSSION

The Organization of the A Domain. The recombinant A domain encompasses the motif of the binding domain truncated, M19 [which includes a trench where, in modeling studies, a triple-helical peptide mimicking collagen was found to fit (13)]. However, the full-length A domain of CNA expressed as a recombinant binds collagen with a higher apparent affinity and specificity: fewer high-affinity sites are found for A compared to M19, and the Kᵦ with the high-affinity binding sites of these two species are 0.3 and 3.0 μM, respectively (12). These results may suggest a model where amino acid residues outside the sequence encompassing M19 continue the structural fold determined by M19. Perhaps the collagen-binding trench in the native protein is extended and involves residues outside the sequence encompassed by M19. An alternative model would suggest that the A domain is composed of subunits that fold independently but may interact to make up the structure of this protein. The presence of an intermediate state in the complex denaturation of the A domain (Figure 5) may support this suggestion, since subunits having different stabilities with respect to heat could yield such a unfolding profile. In this second model, M19 could represent such a subunit. At the present time, we cannot differentiate between these two models.

A and B Domains Behave Independently. Analytical ultracentrifugation measurements demonstrated that the AB₁ heterodimer is more spherical than the A domain, suggesting that the two subunits do not have simply end-to-end joining. To determine whether this behavior is due to reorganization of the two subunits upon joining, we compared the far-UV CD and emission spectra, thermal denaturation profiles, and collagen-binding capabilities of A and AB₁.

Theoretical far-UV CD and tryptophyl emission spectra of AB₁, calculated using eqs 2 and 3, are shown in Figures 9b and 6b. These spectra closely match those obtained experimentally; hence, neither the secondary nor the tertiary structural elements within the A and B₁ subunits undergo detectable reorganization when the two are joined.

Is the complex thermal denaturation observed for AB₁ unique to the heterodimer, a consequence of an interaction between the two domains, or merely a composite of A and B₁ unfolding simultaneously? If the denaturation observed for AB₁ is due only to a sum of the individual subunits (A and B₁) unfolding independently, we should be able to model the curve for AB₁ in Figure 5 by averaging the denaturation curves of A and B₁. The calculated and observed denaturation curves overlay (data not shown), indicating that (1) each subunit unfolds independently and (2) no changes in stability occur as the heterodimer is formed.

This comparison of theoretical and measured profiles of the far-UV CD, emission, and denaturation confirms that the overlap between A and B₁ in the heterodimer does not alter the structure or stability of the two subunits. These results, in conjunction with the collagen-binding measurements shown in Figures 8–10, suggest that the CNA A and B domains do not interact. A plausible construction of the CNA MSCRAMM is shown in Figure 11. In this arrangement, the AB₁ heterodimer would be more globular than the A domain alone, and the two domains make little, if any, contact.

A Model for the CNA B Domain. The far-UV CD spectra of B₁ and B₁B₂ are very similar, indicating that little, if any, alteration in the secondary structure of the individual repeat units occurs upon formation of the B₁B₂ dimer. Not only do B₁ and B₁B₂ have the same percentages of α-helix and β-sheet structural motifs as resolved by a variety of spectral deconvolution programs (Table 3), but also their far-UV CD spectra are almost identical (Figure 4a). The loss of secondary structural organization upon heating occurs similarly for the monomeric and dimeric B domain proteins. No intermediate states in the denaturation were observed for either protein, and the temperature ranges over which the two proteins unfold were alike (Figure 5). A slight increase in T_m is apparent in the dimer, perhaps due to the enhanced

FIGURE 10: Inhibition of recombinant CNA AB₁ to immobilized type I collagen by biotinylated recombinant CNA A. The biotinylated and unbiotinylated species were allowed to bind simultaneously to the immobilized collagen. The dark circles represent self-inhibition (i.e., inhibition of A binding to immobilized type I collagen by biotinylated A), measured as a control. The data points represent average values of duplicate assays.

FIGURE 11: Cartoon drawing of the proposed organization of S. aureus CNA anchored to the cell surface. For simplicity, only the A and B domains are shown.
stability arising from hydrogen bonds and/or the interaction of hydrophobic surfaces at the B$_1$–B$_2$ interface (42).

It is unlikely that the difference in $T_m$ for B$_1$B$_2$ compared to B$_1$ is due to an increased hydrophobicity of tryptophans at the B$_1$–B$_2$ interface, however. From a comparison of emission spectra (Figure 6a), it is evident that the overall tryptophyl environments of B$_1$ and B$_1$B$_2$ are quite similar. No obvious rearrangement of subunit tertiary structure occurs nor do any of the five tryptophans that may be surface-exposed in a single B repeat unit become buried in the interface as two B repeat units are joined.

Forming the B$_1$B$_2$ homodimer causes an increase in the sedimentation velocity frictional ratios (Table 2), implying that the axial ratios and frictional ratio of B$_1$B$_2$ are less than those of B$_1$. This observation suggests that the interface is not a simple end-to-end stacking of the repeat units. The time-resolved fluorescence anisotropy measurements provide supporting evidence for the more globular nature of B$_1$B$_2$ compared to B$_1$. The anisotropy decay is longer-lived for B$_1$B$_2$ than B$_1$, but not tremendously so. If the two repeat units stacked end-to-end, making the dimer a significantly more elongated molecule, we would expect the anisotropy decay of the dimer to be even longer-lived. We propose that the B domain repeat units in CNA join as depicted in Figure 11, where at each unit juncture the protein folds somewhat back onto itself. Since no structural or functional interaction is observed between the repeat units of B$_1$B$_2$, we conclude that there is little significant contact among any of the B domain repeat units.

That (1) no observable reorganization of the secondary or tertiary structure occurs and (2) no dramatic differences upon denaturation are detected upon forming the B$_1$B$_2$ dimer confirms that the two repeat units of B$_1$B$_2$ fold independently and in the same manner as the single B$_1$ unit does. We propose that this autonomy is preserved regardless of the number of repeat units present within a particular strain of CNA-containing S. aureus. Consequently, we limited the mixed-domain analyses to the AB$_1$ recombinant and suggest that this construct is representative of all constructs that include the A domain and any number of B repeat units. Since the A and B domains (and the repeat units within the B domain) are independent, extrapolation of characteristics of the CNA recombinants in vitro to those of the S. aureus MSCRAMM in vivo is possible. It must be kept in mind that if domain cooperativity in protein folding, stability, or collagen binding had been observed, this would not be the case.

No ligand binding has yet been identified for the B domain repeat units, and a B domain is apparently not necessary for presenting a functional binding A domain on the surface of S. aureus cells. Furthermore, the studies presented here suggest that the B domain does not influence the collagen-binding activity of the A domain and the function of the B domain remains unclear.

CNA May Be a Model for Other MSCRAMM Proteins. Surface proteins on Gram-positive bacteria are often composed of unique and repeated sequences. The repeated sequences in a particular protein can occur in different copy numbers depending on the strain of origin. For example, the B domain in CNA may contain 1, 2, 3, or 4 repeats of a 23 kDa unit in the protein from different isolates. This structural organization, which is derived from sequence analysis, suggests that the MSCRAMMs are mosaic proteins. In this study, we demonstrate that CNA is an example of a protein in which genetic motifs translate into individual structural and functional domains in the protein. Furthermore, the CNA A and B domains are independent structurally and functionally, as are the repeat units within the B domain. Thus, CNA is a true mosaic protein. We suggest that these characteristics may be valid for MSCRAMMS in general. Bottomley et al. concluded from their studies of S. aureus protein A, SpA, that the joining the domains in SpA does not affect the structure of the individual domains (43). Also, preliminary analyses of the relationship between domains within other MSCRAMMs support our hypothesis (unpublished results).

The occurrence of different numbers of repeat units in the same protein expressed by nonrelated isolates may have arisen from intragenic recombination, where homologous sequences on each side of the repeat unit represent target sites for crossover. Intronless splicing has recently been suggested to be important in the evolution of prokaryotic protein families. These recer sequences (recombination spacer) are located as structureless links between domains and represent target sites for recombination events. Björck and co-workers described a specific recer sequence presumably involved in the shuffling of modules in surface proteins among Peptostreptococcus magnus strains (44, 45). The recer sequence identified in P. magnus is not found in S. aureus FDA 574 cna, but perhaps other recer classes have evolved among S. aureus strains. Future studies will undoubtedly address these possibilities.

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